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# ***U.S. PATENT APPLICATION***

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***Invention:*** USE OF SYNTHETIC PEPTIDES TO INDUCE TOLERANCE TO  
PATHOGENIC T AND B CELL EPITOPES OF AUTOANTIGENS OR  
INFECTIOUS AGENTS

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## ***SPECIFICATION***

USE OF SYNTHETIC PEPTIDES TO INDUCE TOLERANCE  
TO PATHOGENIC T AND B CELL EPITOPES OF  
AUTOANTIGENS OR INFECTIOUS AGENTS

BACKGROUND OF THE INVENTION

5                   This is a continuation-in-part of  
Application Number 07/833,429, filed February 10,  
1992, which is a continuation-in-part of  
Application No. 07/591,109, filed October 1, 1990,  
which is a continuation-in-part of Application  
10   Number 93,854, filed September 8, 1987, now U.S.  
Patent 5,019,387, the entire contents of which are  
hereby incorporated by reference.

Field of the Invention

15                   The present invention relates, in  
general, to the use of synthetic peptides to  
induce tolerance to immunogenic peptides. In  
particular, the present invention relates to a  
method of inducing tolerance in a mammal to an  
immunogenic peptide or protein comprising  
20   administering to a mammal a synthetic toleragen  
comprising a 2 to 20 amino acid hydrophobic  
peptide linked to the N-terminus or C-terminus of  
the immunogenic peptide or protein, under  
conditions such that the tolerance is induced.

25                   Background Information

Many autoimmune diseases in animals and  
man are characterized by T and B cell responses to  
pathogenic epitopes on self antigens  
(Immunotherapy of Diabetes and Selected Autoimmune  
30   Diseases, G.S. Eisenbarth (ED) CRC Press, Boca  
Raton, 1989; Current Therapy in Allergy,

Immunology, and Rheumatology-3. L.M.

Lichtenstein, et al, B.C. Decker, Inc., Toronto, 1988). Examples of autoimmune diseases or disease models that are caused by autoreactive B cells responses are listed in Table 1. Examples of autoimmune diseases or disease models that are caused by autoreactive T cell responses are listed in Table 2. A method to tolerize human lymphocytes to not respond to pathogenic T and B cell epitopes of autoantigens that otherwise induce immune responses that cause tissue damage would represent a significant advance in the therapy of autoimmune diseases. Similarly, multiple clinical situations exist outside the setting of autoimmune disease, in which B and T cell responses are harmful and would advantageously be shut off or decreased. Examples of pathogenic non-autoimmune antibody responses are antibody responses to ABO incompatible erythrocytes. A method to induce tolerance against this type of immunogen would be a powerful tool for treatment of a number of similar conditions.

Recently, it has also become clear that tissue destruction in certain infectious diseases is caused by immune responses against normal tissue that are induced by infectious agents. For example, in HTLV-I infection, the clinical syndrome of HTLV-1 associated myelopathy (HAM) has been shown to be associated with the induction of cytotoxic T lymphocytes reactive with a specific region (SP4A1) OF HTLV-1 gp46 envelope glycoprotein (S. Jacobson, et al, J. Immunol. 146:1155-1162, 1991). Similarly, lymphocytic pneumonitis in HIV infection has been shown to be associated with the presence in lung lymphocytes of CTL specific for HIV infected cells (AIDS,

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B.D. Walker, et al, 4:177, 1990). In both HIV and HTLV-1 infection, it is thought that certain manifestations of the disease are caused by the induction of anti-viral immune responses that cross-react with normal human host antigens (G.W. Hoffman, et al, Proc. Natl. Acad. Sci. USA 88:3060-3064, 1991; H. Wigzell, et al, FASEB J. p.2406-2410, 1991; H. Golding, et al, J. Clin. Invest. 83:1430-1435).

Robinson et al have demonstrated that antibody responses to HIV envelope gp41 epitopes enhance HIV infectivity (W.E. Robinson, et al, Proc. Natl. Acad. Sci. USA, 86:4710, 1989). Recently, evidence has been presented that many if not all of the manifestations of AIDS may be caused by an autoimmune response to HLA antigens that are induced by HIV viral proteins that share sequence homologies with normal host HLA molecules (G.W. Hoffman, et al, Proc. Natl. Acad. Sci. USA 88:3060-3064, 1991; H. Wigzell, et al, FASEB J. p.2406-2410, 1991; H. Golding, et al, J. Clin. Invest. 83:1430-1435). Thus, a method of induction of tolerance (non-responsiveness) to pathogenic HIV or HTLV-1 protein epitopes (or to epitopes of any other infectious agent that induces autoreactive immune responses), would be an important and novel mode of preventing infectious tissue damage.

The ability to induce tolerance to an immune response induced by an infectious agent to prevent tissue destruction has been proposed as a method of treatment of Herpes simplex virus (HSV-1) corneal inflammation (R.L. Hendricks, et al, J. Immunol. 142:263-269, 1989).

The form of antigen has been suggested to be important regarding determination of whether a protein antigen is an immunogen or a toleragen

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(Reviewed in Weigle (1989) The role of the physical state of human gamma globulin in the in vivo and in vitro induction of immunological tolerance. Chapter 5G, Vol. II, p 51-57). Whereas high molecular weight aggregated gamma globulin is a potent immunogen, low molecular weight globulin is a toleragen (W.O. Weigle, Chpt. 5G, Vol. II, p.51-57, 1989). In this case, the ability of aggregated gamma globulin to induce endogenous IL1 has been suggested as the mode of immunogenicity of aggregated gamma globulin (L.C. Gahring, et al, J. Immunol. 145:1318-1323, 1990)

Others have suggested that some T cell epitopes are inherently immunogenic and some are toleragenic (D.R. Milich, et al, J. Immunol. 143:3148-3156, 1989). Milich has converted toleragenic epitopes of Hepatitis B core antigen to immunogenic epitopes by single amino acid substitutions in the T cell epitopes (D.R. Milich, et al, J. Immunol. 143:3148-3156, 1989). Benacerraf has suggested that freely diffusible antigens are toleragens whereas particulate antigens that are concentrated in cells of the reticuloendothelial system are immunogenic (Benacerraf, B. Properties of antigens in relation to responsiveness and non-responsiveness, in Immunological Tolerance, M. Landy, W. Braun, Eds. Academic Press, NY, NY 1969). In contrast, Nossal reported that the particulate polymeric antigen flagellin was a potent toleragen, and induced tolerance to antibody responses to the Salmonella flagella when injected into neonatal rats (Nossal, G Antigen Dosage in Relation to Responsiveness and Non-responsiveness, in Immunological Tolerance, M. Landy, W. Braun, Eds. Academic Press, NY, NY 1969). Finally, immunogenicity versus toleragenicity of antigens has been proposed to be

due to their affinity of binding to MHC and TCR molecules (rev. in Spent et al, Science 248:1357-2363, 1990).

5 The present invention provides a method  
of modification of peptide immunogens whereby the  
modification changes a potent immunogen into a  
potent toleragen. The invention is based on the  
unexpected observation that the F-domain of HIV-1  
gp41 confers to an antigen the ability to be a  
10 toleragen. Specifically, the hydrophobic N-  
terminal 12 amino acids of the gp41 envelope  
protein that mediate fusion of HIV to uninfected  
cells, the fusogenic (F) domain (M.L. Bosch, et  
al, Science, 244:694-697, 1989), were added C-  
15 terminal to the highly immunogenic T1-SP10 and T1-  
SP10(A) peptides (Table 3) (T.J. Palker, et al, J.  
Immunol. 142:3612-3619; M.K. Hart, et al, J.  
Immunol. 145:2677-2685, 1990; M.K. Hart, et al,  
Proc. Natl. Acad. Sci. USA 88:9448-9452, 1991).  
20 When used as an immunogen in chimpanzees, the T1-  
SP10IIIB and the T1-SP10IIIB(A) peptides were  
potent immunogens, whereas the F-T1-SP10IIIB(A)  
peptide (M.K. Hart, et al, Proc. Natl. Acad. Sci.  
USA 88:9448-9452, 1991) were not as immunogenic at  
25 either low (.1mg/kg) or at high (.5 mg/kg) doses.  
Moreover, challenge of the animals with the highly  
immunogenic T1-SP10IIIB(A) peptide at month 16 of  
the immunization schedule proved that the F-T1-  
SP10IIIB(A) immunized animals were tolerant to the  
30 T1-SP10IIIB(A) HIV gp120 env determinants.

#### SUMMARY OF THE INVENTION

It is a general object of this invention  
to provide a method of inducing tolerance in a  
mammal to an immunogenic peptide or protein.

It is a specific object of this invention to provide a method of inducing tolerance in a mammal to an immunogenic peptide or protein comprising administering to the mammal a synthetic toleragen comprising a hydrophobic peptide linked to the N-terminus or C-terminus of the immunogenic peptide or protein, under conditions such that the tolerance is induced.

Further objects and advantages of the present invention will be clear from the description that follows.

#### BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1. Antibody Titers in ELISA Assay Against Immunizing Peptide Over Time In Chimpanzees Immunized with HIV Env Synthetic Peptides.

Figure 2. Peripheral Blood Mononuclear Cell Proliferative Responses to the T1-SP10IIIIB(A) Peptide in 7 Day Tritiated Thymidine Incorporation Assays.

Figure 3. PBMC Proliferative Responses of Chimpanzees Immunized with T1-SP10 Peptides and F-T1-SP10 Peptides to PHA.

Figure 4. Elution Profile of SP10MN(A) Over a G-75 Sephadex Column.

Figure 5. Elution of T1-SP10MN(A) Over a G-75 Sephadex Column.

Figure 6. Elution of F-T1-SP10MN(A) Over a Sephadex G-75 Column.

Figure 7. Elution of F-SP10MN(A) Over G-75 Column.

Figure 8. Results of DSP Cross-linking Analysis Using F-T1-SP10IIIIB(A) Peptide.

Figure 9. Hypothetical Model of F-T1-SP10IIIIB(A) in Aqueous Solution.

Figure 10. Variants of T1-SP10 peptides derived from HIV MN and IIIB Envelope Sequences.

Figure 11. Time course of PBMC 3H-thymidine incorporation responses to HIV Th-B peptide, T1-SP10IIIB(A), in chimpanzees immunized with HIV envelope synthetic peptides. Animals 884 (Figure 1A) and 1028 (Figure 1B) received the Th-B peptide, T1-SP10IIIB, initially (months 1-5), then the Th-B peptide, T-SP10IIIB(A) (month 6-8). After a boost with the Th-B peptide T-SP10IIIB(A) at month 14, both animals 884 and 1028 were immunized with the HIVMN Th-B peptide, T-SP10MN(A). Panels C and D show the responses of animal 1045 (Panel C) and 1070 (Panel D) to the HIVIIIB F-Th-B peptide (month 1-14), HIVIIIB Th-B peptide (month 16) and HIVMN Th-B peptide (months 17-19). All immunizations were with the indicated peptide in IFA, except all immunizations for animal 1028 after month 4, which were with peptides in PBS alone. Solid lines show data for peak proliferative responses ( $\Delta$ cpm) to a wide dose range of HIVIIIB Th-B peptide. Dotted lines indicate peak proliferative response ( $\Delta$ cpm) to a wide dose range of the HIVMN Th-B peptide.

Figure 12. Time course of PBMC 3H-thymidine incorporation response to PHA in chimpanzees immunized with HIV envelope synthetic peptides. Immunizations and chimpanzees as in Figure 11.

Figure 13. Time course of PBMC 3H-thymidine incorporation response to candida antigen in chimpanzees immunized with HIV envelope synthetic peptides. Immunizations and chimpanzees as in Figure 11.

Figure 14. Time course of absolute numbers of lymphocytes and lymphocytic subsets in chimpanzees immunized with HIV envelope synthetic

peptides. Immunizations and chimpanzees as in Figure 11. Points represent cell number/mm<sup>3</sup> of peripheral blood lymphocytes and lymphocyte subsets. The elevated cell numbers in animal 1028 at month 4 coincided with an abscess at the injection sites.

Figure 15. HIV envelope hybrid synthetic peptides induced anti-HIV neutralizing antibodies in goats. Goat 102A was immunized with 3 mg of the F-Th-B peptide, F-T1-SP10IIIIB(A) and goat 104A was immunized with the HIVIIIIB Th-B peptide, T1-SP10IIIIB. Immunizations were in CFA (first dose) and IFA (doses 2-4). Neutralizing titers are titers at which reverse transcriptase production was inhibited by 90% or greater.

#### DETAILED DESCRIPTION OF THE INVENTION

The present invention relates to a procedure whereby protein immunogens are derivatized by either synthesizing a hydrophobic amino acid sequence of 2 to 20 amino acids in length, N-terminal to the immunogenic protein or protein fragment, or covalently linking a hydrophobic peptide fragment of 2 to 20 amino acids in length N-terminal to the immunogenic protein or protein fragment, to yield an immunogen capable of inducing tolerance to the protein immunogen when administered to a mammal such as a primate (more preferably, humans).

In a preferred embodiment, the hydrophobic peptide is 5 to 15 amino acids in length. In yet another preferred embodiment, the hydrophobic peptide is 7 to 13 amino acids in length. In a further embodiment, the length of the hydrophobic peptide is 7, 8, 9, 10, 11, 12, or 13 amino acids in length. In yet another

embodiment, the hydrophobic peptide is at least 5 amino acids in length (more preferably, at least 10 amino acids in length).

Alternatively, immunogenic proteins known to be the targets of autoantibody or auto-T cell responses can be constructed using recombinant DNA technology to form new toleragens containing hydrophobic N-terminal regions as described above. While an advantageous construction of the invention is for the hydrophobic sequence to be N-terminal to the B or T cell epitope, in certain circumstances it may be advantageous to have the hydrophobic sequence C-terminal to the B or T cell epitope.

The hydrophobic region can be a fusion protein from HIV or HIV-related viruses (see Table 5), or can be a hydrophobic sequence of amino acids that is either randomly selected or is from a non-HIV related protein.

An example of this invention for inducing tolerance to antibodies against autoantigens is for the treatment of myasthenia gravis, whereby the F-sequence is synthesized N-terminal to the main immunogenic region of the acetylcholine receptor, **WNPADYGGIK** or **WNPDDYGGVK** (I. Papdoulis, et al, Biochem. J. 269:239-245, 1990). The resulting immunogen is **AVGIGALFLGFLWNPADYGGIK** or **AVGIGALFLGFLWNPDDYGGVK**.

Another example of a B cell toleragen is a hybrid protein comprising the HIV fusion domain synthesized either linearly N-terminal to B cell peptide epitopes of the insulin molecule or covalently linked to the whole insulin molecule or covalently linked or constructed using recombinant DNA techniques to a peptide insulin fragment or to the whole insulin molecule. The resulting immunogen is **AVGIGALFLGFL-insulin** or **AVGIGALFLGFL-**

insulin peptide fragment. These types of  
toleragens can be used to prevent the onset of  
juvenile diabetes mellitus, (J.P. Palmer, et al,  
Science 222:1337-1339, 1983; B.M. Dean, et al,  
5 Diabetologia 23:339-342, 1986) and to treat  
patients with insulin antibodies in the setting of  
insulin resulin resistance (J.D. Schnatz,  
Dolovich, et al, J. Allergy, 46:127-1137, 1970).

Another example of a B cell toleragen is  
10 a hybrid protein comprising the HIV fusion domain  
synthesized either linearly N-terminal to B cell  
peptide epitopes of the TSH receptor molecule or  
covalently linked to the whole TSH receptor  
molecule or covalently linked or constructed using  
15 recombinant DNA techniques to a peptide TSH  
receptor fragment or to the whole TSH receptor  
molecule. The resulting immunogen is  
AVGIGALFLGFL-TSH receptor or AVGIGALFLGFL-TSH  
receptor peptide fragment. These types of  
20 toleragens can be used to treat autoimmune thyroid  
disease (Graves' Disease) (T. Mori, et al,  
Biochem. & Biophy. Res. Comm. 178:165-172, 1991;  
M. Murakami, et al, Biochem. & Biophy. Res. Comm.  
171:512-518, 1990). Table 6 summarizes B cell  
25 epitopes on the thyrotropin (TSH) receptor to  
which Graves' patient sera bind (T. Mori, et al,  
Biochem. & Biophy. Res. Comm. 178:165-172, 1991;  
M. Murakami, et al, Biochem. & Biophy. Res. Comm.  
171:512-518, 1990; O. Takai, et al, Biochem. &  
30 Biophy. Res. Comm. 179:319-326, 1991;  
T. Piraphatdis, et al, Biochem. & Biophy. Res.  
Comm. 172:529-536, 1990). Of interest is the  
sequence YYVFFEEQEDEIIGF identified by 2 studies  
that inhibits the TSH activity of the  
35 autoantibodies (T. Mori, et al, Biochem. & Biophy.  
Res. Comm. 178:165-172, 1991; O. Takai, et al,  
Biochem. & Biophy. Res. Comm. 179:319-326, 1991).

Thus constructs for inducing tolerance to anti-TSH antibodies in Graves' disease are

AVGIGALFLGFLYVVFEEQEDEI or

AVGIGALFLGFLHQEEDFRVTCKDIQRIPSLPPSTQT or

5 AVGIGALFLGFLLRQKSVNALNSPLHQEYBENLGDSIVGY or

AVGIGALFLGFLYYVVFEEQEDEIIGF or

AVGIGALFLGFLYKELPLLKFL.

10 An example of the use of this invention in the induction of tolerance to autoimmune T cell antigens is a hybrid protein comprised of the HIV fusion domain synthesized either linearly N-terminal to T cell peptide epitopes of the myelin basic protein molecule or covalently linked or constructed using recombinant DNA techniques to a  
15 myelin protein molecule. The resulting immunogen is AVGIGALFLGFL-myelin basic protein or AVGIGALFLGFL-myelin basic protein peptide fragment. In the case of the myelin basic protein peptide fragment, the encephalitogenic T cell  
20 epitopes are known, one of which is contained in sequence 69-89 of bovine myelin basic protein (H. Offner, et al, J. Immunol. 141:3828-3832, 1988). In this case, one formulation of the toleragen is AVGIGALFLGFLGSLPQKSQRSQDENPVVHF. These types of  
25 toleragens can be used to treat experimental autoimmune encephalomyelitis, which is thought to be an excellent model of human multiple sclerosis (K.W. Wucherpfenning, et al, Immunol. Today, 12:277-281, 1991). When the specific epitopes are  
30 identified that are the T cell targets in multiple sclerosis, then those sequences can be substituted in the peptide above, and used to tolerize T cells to the pathogenic T cell epitope of whatever the protein antigen turns out to be involved in  
35 multiple sclerosis.

Another example of this invention for induction of tolerance to autoimmune T cell

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antigens is a hybrid protein comprising the HIV fusion domain synthesized either linearly N-terminal to T cell peptide epitopes of the retinal S protein molecule or covalently linked to the whole retinal S protein molecule or covalently linked or constructed using recombinant DNA techniques to retinal S antigen fragment or to the whole retinal S antigen molecule. The resulting immunogen is **AVGIGALFLGFL-retinal S protein** or **AVGIGALFLGFL-retinal S protein peptide fragment**. In the case of the retinal S protein peptide fragment, the pathogenic T cell epitopes are known, one of which is present in the sequence 1169-1191 of retinal S protein (H. Sanui, et al, Exp. Med., 169:1947-1960, 1989). In this case, one formulation of the toleragen is **AVGIGALFLGFLPTARSVGAADGSSWEGVGVV**. These types of toleragens can be used to treat experimental autoimmune retinouveitis, which is thought to be an excellent model of human inflammatory eye diseases such as Bechet's syndrome and idiopathic retinouveitis (H. Sanui, et al, Exp. Med., 169:1947-1960, 1989). When the specific epitopes are discovered that are the T cell targets in human inflammatory eye disease, then those sequences can be substituted in the peptide above, and used to tolerize T cells to the pathogenic T cell epitope of whatever the protein antigen turns out to be in human retinouveitis.

For the treatment of pathogenic immune responses induced by an infectious agent, an example of the invention is the treatment of HTLV-I associated myelopathy syndrome seen in tropical spastic paraparesis (rev. in Jacobson et al J. Immunol. 146:1155-1162, 1991). In this disease, there is strong evidence that the neurologic disease is caused by the induction of

cytotoxic T cells (CTL) against HTLV-I infected cells in the central nervous system (S. Jacobson, et al, J. Immunol. 146:1155-1162, 1991).

Jacobson, et al have shown that one primary region  
5 of HTLV-I env gp46 that induces CTL in tropical  
spastic paraparesis (TSP) is aa196-209 of gp46 as  
defined by peptide SP4a1 (S. Jacobson, et al, J.  
Immunol. 146:1155-1162, 1991; T.J. Palker, et al,  
J. Immunol., 142:971-978, 1989; A. Kurata, et al,  
10 J. Immunol., 143:2024-2030, 1989). Thus, to treat  
TSP, the present invention can be embodied by the  
hybrid peptide AVGIGALFLGFLLDHILEPSIPWKS~~KK~~. When  
new pathogenic CTL epitopes of HTLV-I are  
discovered, the therapeutic construct can be F-X  
15 where F is the hydrophobic sequence and X is the  
CTL epitope of the infectious agent.

The clinical manifestations of HIV have  
been postulated to be due to autoimmune responses  
induced by components of HIV that have sequence  
20 homology to human MHC Class I or Class II  
molecules (G.W. Hoffman, et al, Prac. Natl. Acad.  
Sci. USA 88:3060-3064, 1991; H. Wigzell, et al,  
FASEB J. p.2406-2410, 1991; H. Golding, et al,  
J. Clin. Invest. 83:1430-1435; F. Grassi, et al,  
25 J. Ex. Med., 174:53-62, 1991; J.A.T. Young,  
Nature, 333:215, 1988; H. Golding, et al, J. Exp.  
Med., 167:914-923, 1988). For the treatment of  
HIV infection, the present invention can comprise  
a series of hybrid peptides, each peptide  
30 containing an N-terminal hydrophobic peptide such  
as the HIV gp41 fusion domain (Table 5) and a C-  
terminal peptide from each of the regions of HIV  
env proteins bearing sequence homology MHC class I  
or class II molecules (G.W. Hoffman, et al, Prac.  
35 Natl. Acad. Sci. USA 88:3060-3064, 1991; H.  
Wigzell, et al, FASEB J. p.2406-2410, 1991; H.  
Golding, et al, J. Clin. Invest. 83:1430-1435; F.

Grassi, et al, J. Ex. Med., 174:53-62, 1991;  
J.A.T. Young, Nature, 333:215, 1988; H. Golding,  
et al, J. Exp. Med., 167:914-923, 1988) (Table 7).  
Alternatively, it may be advantageous to treat HIV  
5 infected individuals with F-X peptides where F is  
a hydrophobic peptide such as the fusogenic domain  
of HIV and X is a peptide fragment of HIV that is  
immunogenic to T or B cells. In this situation, a  
mixture of peptides would be used to inhibit  
10 destructive anti-HIV immune responses that were  
damaging host HIV-infected antigen-presenting  
cells. Examples of this type of peptide are shown  
in Table 3 and Figure 10, and were the peptides  
used that tolerized chimpanzees in Figures 1 and 2  
15 to both T1-SP10(A) determinants and to whole gp120  
protein (Table 4).

The present invention is described in  
further detail in the following non-limiting  
Example.

#### 20 Example 1

Antibody titers in ELISA assay against  
immunizing peptide over time in chimpanzees  
immunized with HIV env synthetic peptides are  
shown in Figure 1. For animals 884 and 1028, the  
25 peptide used in the ELISA assay was T1-SP10IIIB.  
For animals 1045 and 1070, the peptide used in the  
ELISA assay was F-T1-SP10IIIB(A). All  
immunizations were in incomplete Freund's Adjuvant  
(IFA) + PBS (1:1) except for animal 1028 that  
30 developed IM abscesses after immunization no. 3,  
and had one immunization held, then had all  
subsequent immunizations in PBS only. As can be  
seen, T1-SP10 peptides were excellent immunogens  
in animals 884 and 1028, while T1-SP10 peptides  
35 with the HIV gp41 fusion (F) domain synthesized N-

terminal to the T1-SP10 peptide did not induce antibody titers as high or as of long duration as did peptides without the F domain.

It is important to note that animals 1045 and 1070 were challenged at month 16 with the immunogen T1-SP10IIIB(A) that induced such good antibody titers in animals 884 and 1028. Animals 1045 and 1028 did not respond to T1-SP10IIIB(A) in IFA, thus demonstrating that they were tolerant to the T1-SP10(A) from their prior immunizations with F-T1-SP10IIIB(A) peptide. It is also important to note that while boost of 884 at week 14 gave a rise in titer to T1-SP10IIIB(A) peptide, boost of 1028 at the same time did not. Boost of 884 was with IFA, while boost of 1028 was with no adjuvant, but rather only PBS.

Peripheral blood mononuclear cell proliferative responses to the T1-SP10IIIB(A) peptide in 7 day tritiated thymidine incorporation assays is shown in Figure 2. T1-SP10IIIB and T1-SP10IIIB(A) peptides induced high levels of proliferation of circulating PBMC in animals 884 and 1028. These levels fell to non-detectable levels after a 6 month rest (month 14) but rose again in animals 884 and 1028. Proliferative responses in animal 1028 rose with each boost after the 6 month rest even though the immunizations were in PBS alone with no adjuvant. As with B cell response, animals 1045 and 1070, immunized with F-T1-SP10IIIB(A) peptide, did not proliferate to T1-SP10IIIB(A) peptide. When these latter two animals were immunized with the T1-SP10IIIB(A) peptide that was a good immunogen in 884 and 1028, neither of the animals 1045, 1070 developed a proliferative response to T1-SP10IIIB(A)--thus proving that the addition of the F-domain N-terminal to the T1-SP10 peptide created

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a toleragen that tolerized animals 1045 and 1070 to the T1 and SP10 regions of gp120. As shown in Table 4, while animals 884 and 1028 both responded in proliferative assays to native gp120, animals 1045 and 1070 were tolerant to native gp120 as well as to immunizing peptides.

PBMC proliferative responses of chimpanzees immunized with T1-SP10 peptides and F-T1-SP10 peptides to PHA are shown in Figure 3.

Data show that while animals 1045 and 1070 were tolerant to T1 and SP10 regions of HIV gp120, PBMC PHA responses in these animals throughout the immunization period were normal.

Similar results were obtained with peripheral blood mononuclear cell (PBMC) responses to candida antigen in 7 day in vitro stimulation assay (not shown). Thus, while specifically tolerant to T1 AND SP10(A), HIV env determinants, animal 1029 and 1045 were not generally immunosuppressed and could respond to candida to PHA stimulation in vitro.

The effect on peptide quaternary structure of placement of a hydrophobic sequence N-terminal to a T cell and/or a B cell determinant was examined. Using G-75 chromatography in aqueous buffers and crosslinking of peptide monomers using the heterobifunctional agent Dithiobis (succinimidylpropionate) (DSP), it was determined that addition of a hydrophobic sequence such as the fusion (F) domain of HIV or HIV-like retroviruses confers on the T1-SP10(A) or the SP10(A) peptide the ability to form high molecular weight aggregates, that are likely in the form of protein micelles.

An elution profile of SP10MN(A) over a G-75 Sephadex column is shown in Figure 4. 4mg of each peptide in 2ml 50mM Tris-HCl (pH 7.5)

containing 100mM KCl and 5% glycerol, was applied directly to a 90 x 1.6 cm column of Sephadex G-75 equilibrated with 50mM Tris-HCl (pH 7.5) containing 100mM KCl. The sizing column was calibrated with blue dextran (200,000), bovine serum albumin (66,000), bovine erythrocyte carbonic anhydrase (29,000), horse heart cytochrome C (12,400) and bovine lung aprotinin (6,500). The elution position of each peptide was determined by continuous measurement of eluent absorbance at OD 280. The corresponding molecular weight of each peptide peak was calculated from the calibration curve of the column. Each peptide was also applied to the column equilibrated with the same buffer containing 0.1% C12E9 [polyoxyethylene (9) lauryl ether]. The SP10MN(A) peptide (predicted Mr=2878) migrated as forms of 65 Da or lower. Similarly, the T1-SP10MN(A) (predicted Mr=4771) peptide also migrated as low mw forms ranging from 12,000 Da to 6,500 Da (Figure 5). In contrast, both the F-SP10MN(A) (Mr=4038) and the F-T1-SP10MN(A) peptides (Mr=5930) contained high molecular weight forms that migrated at ~66,000 Da (Figures 6 and 7). Methods used in Figures 5, 6, and 7 were as in Figure 4.

The results of DSP cross-linking analysis using F-T1-SP10IIIIB(A) peptide are shown in Figure 8. Lane C shows the form of the peptide with no DSP added in PBS when run under non-reducing conditions in SDS-PAGE. Lanes D,E,F, and G show the effect on peptide MW when the peptide is cross-linked with 6.25 $\mu$ g (D), 12.5 $\mu$ g (E), 25 $\mu$ g (F) and 50 $\mu$ g (G) of DSP prior to SDS-PAGE. Lane H shows the results of addition of 2-ME to peptide cross-linked with 50 $\mu$ g of DSP and then run under reducing conditions in SDS-PAGE showing all of the

cross-linked forms seen in lane H and all the multiple forms seen in non-reduced, non-cross-linked peptide seen in lane C, were now reduced to two bands at 7000 kDa. At present, the nature of the two bands in this peptide under reducing conditions is unknown; these two bands can be purified by cutting the bands out of preparative gels and can be analyzed by mass spectroscopy and sequenced. Lanes A and B show the results of crosslinking F-T1-SP10IIIB(A) peptide in the presence to Triton-X 100 1% and run under reducing (A) and non-reducing conditions (B). Data demonstrate that the apparent hydrophobic interactions holding the high MW complexes together are resistant to disruption by this detergent.

A hypothetical model of F-T1-SP10IIIB(A) in aqueous solution is shown in Figure 9. The model shows protein micelle formation with the hydrophobic fusion domain (F) regions of the peptide in the core of the micelle with the hydrophilic V3 regions projecting outward.

#### Examples 2-8

The following experimental details and protocols are referenced in Examples 2-8.

Peptides: Peptides used in Examples 2-8 that follow are listed in Table 8. Peptide synthesis was performed using either t-boc or f-moc chemistry with a peptide synthesizer (A431; Applied Biosystems, Inc. Foster City, CA). Peptides were purified using HPLC, and the molecular weight was determined by fast atom bombardment mass spectrometry (R. B. Van Breeman, North Carolina State University, Raleigh, NC) using a double-focusing mass spectrometer

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(HXIIIOHF; Joel Ltd., Tokyo, Japan). For Th-B and F-Th-B peptides (Table 8), expected molecular mass of F-Th-B peptide, F-T1-SP10IIIB(A), was 5908, observed was 5907; expected molecular weight of Th-B peptide, T1-SP10IIIB, was 4061, observed was 4062; expected and observed molecular weight of Th-B peptide, T1-SP10IIIB(a) was 4,749, and expected and observed molecular weight of Th-B peptide, T1-SP10MN(A), was 4771. For the peptides used in the following Examples (Table 8), the peptide amounts are gross weights. The % water by Karl Fisher Test (Galbraith Laboratories, Inc. Knoxville, TN) for each peptide was F-T1-SP10IIIB(A), 6%; T1-SP10IIIB(A), 8%; T1-SP10IIIB, 6% and T1-SP10MN(A), 8%.

Animals: Chimpanzees were housed at the New Mexico State University Primate Facility at Alamogordo, NM. Chimpanzee No. 884 (15 yrs. old) and 1028 (12 yrs. old) had the same sire; animal 1045 (10 yrs. old) and 1070 (11 yrs. old) were unrelated to each other and to animals 884 and 1028. Outbred goats were housed at the Duke University Animal Facilities.

Immunizations: For goats, 3 mg of peptide were injected intramuscularly in each gluteal region in complete Freund's adjuvant (CFA) (1st dose), then incomplete Freund's adjuvant (IFA) (subsequent doses). For immunization of chimpanzees, varying doses of peptides were injected IM in IFA in a total volume of 4 cc, with 1 cc injected into right and left upper arms and thighs.

ELISA Assays: 2  $\mu$ g of Th-B peptide, T1-SP10IIIB, or rgp120IIIB (Repligen Corp., Cambridge, MD) in CBC buffer (15 mM  $\text{Na}_2\text{CO}_3$ , 35 mM  $\text{NaHCO}_3$ , pH9.6) was incubated overnight in each well of a 96 well flat bottom plate (Costar 3590).

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Wells were blocked with CBC buffer supplemented with 3% bovine serum albumin (BSA) for at least 2 hrs and then were washed 3 times with PBS, 0.05% Tween 20. Primary antibody at various concentrations in serum diluent (95 ml PBS, 0.05% Tween 20, supplemented with 5 g BSA in 2 ml normal serum from same species as secondary antibody) was incubated for 90 min at 20°C. After washing three times, alkaline phosphatase-conjugated secondary antibody was added to each well (60 min at RT) and the plates washed. Substrate (1 mg/ml p-nitrophenyl phosphate, Sigma Chemical Co., St. Louis, MO) in 0.05M CBC-0.002M MgCl<sub>2</sub> was added to each well, and plates developed (60 min, 20°C) in the dark and read at 405 nm on an ELISA reader (Anthros; Denley Instruments Co., Durham, NC). Endpoint ELISA antibody titers were defined as the serum titer at which the experimental/control (E/C) OD value  $\geq 3.0$ .

HIV Neutralization Assays: The ability of chimpanzee or goat serum antibodies to neutralize HIV was determined in syncytium inhibition assay and reverse transcriptase inhibition assay as previously described (Palker et al, J. Immunol. 142:3612 (1989); Palker et al, Proc. Natl. Acad. Sci. USA 85:1932 (1988)). Sera were heat inactivated (30 min, 56°C) prior to each assay.

PBMC Isolation and In Vitro <sup>3</sup>H-Thymidine Incorporation Assays: Chimpanzee or goat PBMC was isolated by standard density centrifugation techniques (Palker et al, J. Immunol. 142:3612 (1989); Haynes et al, Science 215:298 (1982)). In vitro assays of <sup>3</sup>H-thymidine incorporation were performed as described (Palker et al, J. Immunol. 142:3612 (1989); Hart et al, J. Immunol. 145:2697 (1990)). For chimpanzee PBMC assays, in vitro

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cultures were performed using 10% normal chimpanzee serum. Antigens used in PBMC proliferation were the Th-B peptides, T1SP10IIIB(A) and T1-SP10MN(A), (Table 8), and Candida albicans antigen (Greer Laboratories, Inc. Lenoir, NC). PHA (Burroughs Wellcome, Research Triangle Park, NC) was used in a wide dose range as a mitogen in 3 day PBMC <sup>3</sup>H-thymidine incorporation assays (Palker et al, J. Immunol. 142:3612 (1989); Hart et al, J. Immunol. 145:2697 (1990)). Acpm = experimental cpm - control cpm.

Immunization Schedule: Because of previous studies demonstrating the immunogenicity of Th-B peptides in goats and rhesus monkeys (Hart et al, J. Immunol. 145:2697 (1990)), the initial comparison of peptide designs when this study began in 1989 was monthly injections of Th-B versus F-Th-B peptides (Table 8) at a dose of approximately 0.1 mg/kg (6 mg/animal). When neither peptide design induced neutralizing anti-HIVIIIB antibodies, the peptide doses were increased to approximately 0.5 mg/kg (30 mg/animal) and the right-hand side neutralizing sequence of HIVIIIB gp120 V3 loop (the (A) region) (Hart et al, Proc. Natl. Acad. Sci. USA 88:9448; Rusche et al Proc. Natl. Acad. Sci. USA 85:3198)) (Table 8) was added to the Th-B peptide to enhance the ability of this peptide to induce anti-HIVIIIB neutralizing antibodies. After 3 monthly injections with either ~0.5 mg/kg (30 mg) Th-B or F-Th-B peptide, the animals were rested for 6 months, and then reimmunized with either F-Th-B or Th-B with sequences from HIVIIIB, or with the Th-B peptide containing HIV env gp120 V3 sequences from the HIVMN isolate.

Flow Cytometry: Chimpanzee PB mononuclear cells were studied by standard flow

cytometry methods using a flow cytometer (751; Coulter Electronics, Inc., Hialeah, FL). PB lymphocytes were identified by the following markers; total T cells, CD3; T cell subunits, CD4 and CD8; B cells, CD19; and NK cells, CD56 and CD16.

## Example 2

### Immunogenicity of Th B and F Th B Peptides in Chimpanzees and Goats for Anti-Peptide and Anti-HIV gp 120 Antibody Responses

For chimpanzees immunized with HIVIIIB Th-B peptides (chimpanzee nos. 884 and 1028), antibody to immunizing peptide rose during the initial immunization period (Table 9). Chimpanzee no. 1028 developed an abscess at the immunization site, did not receive the month 5 immunization, and all subsequent immunizations after month 5 in animal 1028 were in PBS alone. Whereas peak endpoint ELISA anti-peptide antibody titer at month 4 in animal 1028 was 1:819,200, antibody titers fell in animal 1028 after IFA was deleted from the immunogen, and remained low throughout the remainder of the immunization period (Table 9). In chimpanzee no. 884, antibody titers rose at month 7 to 1:204,800 after 5 immunizations with Th-B peptides. Continued immunization of animal 884 with high doses of Th-B peptide (30 mg/dose) resulted in no further increases in antibody titer (Table 9).

In contrast, anti-peptide antibody levels were much lower during months 1-10 of immunization of animals 1045 and 1070 with HIVIIIB F-Th-B peptide, with peak antibody levels against immunizing peptide of 1:25,600 and 1:12,800 at

month 7 for animals 1028 and 1070, respectively (Table 9). After a 6 month rest for all four animals, animals 884 and 1028 were immunized at month 14 with 6 mg of Th-B peptide. In chimpanzee no. 884, boosting with Th-B peptide in IFA at month 14 resulted in rise in titer of anti-peptide antibody to 1:102,400, while boosting of animal 1028 with peptide in PBS alone led to no antibody rise (Table 9).

In contrast, animals 1045 and 1070 were immunized at month 14 with 1 mg (~0.016 mg/kg) of F-Th-B to determine if the prior doses of F-Th-B peptide were excessive and induced high zone tolerance, and if smaller amounts of F-derivatized peptide would be more immunogenic. Immunization of both chimpanzee nos. 1045 and 1070 with 1 mg of F-Th-B peptide after a 6 month rest resulted in only minimal rises in serum titers of anti-peptide antibody to 1:800 (Table 9).

To determine if chimpanzees 1045 and 1070 were tolerant to Th-B peptides, both animals were immunized on month 16 with HIVIIIB Th-B peptide, T1-SP10IIIB(A). Both animals 1045 and 1070 responded minimally to boosting with Th-B peptide with an antipeptide antibody responses to 1:1600 and 1:3200, respectively, demonstrating that animals 1045 and 1070 were hyporesponsive at month 16 to Th-B HIV env epitopes (Table 9).

### Example 3

Immunization of Animals 1045 and 1070 with HIVMN Th-B Peptide Induced High Levels of Antipeptide Antibodies

Using a previously described strategy of breaking B cell tolerance by immunization with an

immunogen that is different from, but structurally related to, the tolerogen (Weigle, Natural and Acquired Immunologic Unresponsiveness (1967) Chapter 4, pp. 57-151), animals 1045 and 1070 were next immunized with the HIVMN Th-B peptide. The TH-B peptide from HIVMN contained the same Th (T1) gp 120 sequence as the HIVIIIB Th-B peptide, but contained different B cell gp 120 V3 B cell epitope sequences than those in the HIVIIIB Th-B peptide (Table 8). After 2 immunizations with Th-B of HIVMN, beginning at month 17, both chimpanzee nos. 1045 and 1070 had prompt rises in titer of antibodies to HIVIIIB (Table 9) and to HIVMN Th-B peptide (not shown) to antibody levels that were higher than had previously been obtained during the prior 18 months of study. At month 20, endpoint ELISA titers to the HIVMN Th-B peptide were 1:102,400 for animal 1045 and 1:204,800 for animal 1070.

#### Example 4

##### Chimpanzee B Cell Antibody Responses to Recombinant HIVIIIB gp120 During the 20-Month Immunization Course

Endpoint ELISA antibody titers against recombinant HIVIIIB gp 120 were determined for sera from months 4-7 and 16-20 to correlate peak anti-peptide antibody levels with anti-gp120 HIV envelope antibody levels. It was found that peak anti-gp120 antibody levels in chimpanzee nos. 884 and 1028 during months 4-7 were both 1:25,600, whereas peak titers to gp120 in animals 1045 and 1070 during the same period were 1:6,400 and 0, respectively. As with anti-peptide antibody levels, boosting after a 6 month rest with peptide

in PBS in chimpanzee 1028 did not boost anti-gp120 antibodies.

Boosting with F-Th-B peptide at month 14 and with HIVIII TH-B at month 16 in animals 1070 and 1045 resulted in minimal rises in anti-gp120 antibody titers by month 17 (to 1:12,800). In contrast, boosting chimpanzees 1045 and 1070 with HIVMN Th-B peptide at month 17 induced high levels of anti-gp120IIIB antibody in both animals (1:102,400 and 1:51,200, respectively) by month 20 that rose coincident with rises in levels of anti-peptide antibody.

#### Example 5

#### Induction of Anti-Peptide and Anti-gp120 PBMC Proliferative Responses by HIV Env Peptides

Whereas HIVIIIIB Th-B peptides induced high levels ( $>100,000$  Acpm/ $10^6$  cells) of PBMC  $^3$ H-thymidine incorporation (animals 884 and 1028) (Figures 11A and 11B) during months 1-8, F-Th-B peptide did not induce levels of  $^3$ H-thymidine incorporation above  $100,000$  Acpm/ $10^6$  cells during the same period (Figures 11C and 11D). Immunization of animals 1045 and 1070 with Th-B peptide at month 16 did not induce the presence of circulating PBMC capable of proliferating to Th-B peptide in vitro (Figures 11C and 11D).

Interestingly, Th-B peptides at month 14-18 boosted PBMC proliferative responses in animal 1028, while anti-peptide antibody responses in animal 1028 during this time were not boosted (Figure 11B and Table 8).

Next,  $^3$ H-thymidine incorporation of chimpanzee PBMC to either recombinant gp120IIIB or

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to native gp120IIIB was tested. Table 4 shows the peak <sup>3</sup>H-thymidine incorporation of chimpanzee PBMC to HIVIIIB gp120 for each animal during months 1-13, and demonstrates that neither chimpanzee no. 1070 nor 1045 (receiving F-Th-B peptide) had PBMC-proliferative responses to gp120 of greater than E/C>2 throughout the first 13 months of study. In contrast, animals 884 and 1028 (receiving Th-B peptides) did have anti-gp120 proliferative responses during the same period (Table 4).

To determine if PBMC proliferative responses to mitogenic or antigenic stimuli other than HIV immunogens were normal in the F-Th-B-immunized chimpanzees over the 20 months of study, we also measured PBMC proliferative responses to PHA were also measured (Figure 12) and to Candida (Figure 13). While peak PHA PBMC proliferative responses were nearly identical in the four chimpanzees, Candida PBMC-proliferative responses varied from animal to animal and from month to month. However, in animals 1045 and 1070, it was found that Candida responses were intermittently present during the time of immunization with F-Th-B peptide at levels that were similar to levels present before the immunizations were begun (Figures 13C and 13D).

#### Example 6

##### Characterization of PB Lymphocyte Subsets During Immunization of Chimpanzees With HIV Env Peptides

To determine if immunization with either HIV env peptide type had effects on the number of circulating chimpanzee T, B or NK cell populations, the absolute numbers of these cell

types were determined throughout the immunization period (Figure 14, Table 10). Whereas preimmunization (before) and postimmunization (during) lymphocyte levels in animals 884 and 1028 were not significantly different (Table 10), animal 1045 became relatively lymphogenic ( $p < 0.001$ ) during the course of immunization with F-Th-B peptide with the lymphocyte count  $650/\text{mm}^3$  at week 12, compared to preimmunization levels of 2815 and 2597 lymphocytes/ $\text{mm}^3$  in months 1 and 2, respectively (Figure 14C). Whereas T cell levels significantly dropped an average of 59% and 44% in chimpanzee nos. 1045 ( $p > 0.001$ ) and 1070 ( $p > 0.02$ ), respectively, during the immunization period, T cell levels did not significantly change in animals 884 and 1028 during the same time ( $p > 0.1$ ) (Table 10). B and NK cell levels dropped significantly in animal 1045, but did not change in animals 1070, 884 and 1028 (Table 10). Taken together, these data demonstrated that immunization with the F-derivatized HIV env peptide induced decreases in absolute levels of circulating T cells in both animals 1045 and 1070, and in B and NK cell levels in animal 1045, whereas immunization of chimpanzee nos. 884 and 1028 with HIV Th-B env peptides lacking the F domain did not significantly affect circulating lymphocyte levels.

#### Example 7

Ability of HIVIIIIB F-Th-B and Th-B Peptides to Induce Anti-HIVIIIIB Neutralizing Antibodies in Goats

To determine if the F-Th-B peptide used in the initial phase of the chimpanzee

immunization protocol was immunogenic in another species, 3 mg of either F-Th-B or Th-B peptide were used to immunize goats three times over 2 months and then used to boost goats after an 8 month rest (Figure 15). It was found that after the fourth immunization, both peptides were capable of inducing serum anti-HIVIIIB neutralizing antibodies (Figure 15), and capable of inducing high levels ( $\geq 500,000$  Acpm/ $120^6$  cells) of PBMC  $^3\text{H}$ -thymidine incorporation in vitro to Th-B or F-Th-B peptides. In addition, serum endpoint ELISA titers of antibodies to immunizing peptide were the same in Th-B and F-Th-B-immunized goats. Thus, failure of the F-Th-B peptide to induce high levels of anti-peptide antibodies and PBMC-proliferative responses in chimpanzees was not due to lack of an inherent immunogenicity of the HIVIIIB F-Th-B peptide, but rather was due to a specific effect of the F-derivatized peptide in chimpanzees.

#### Example 8

##### HIVMN Th-B Env Peptide Induced Anti-HIV Neutralizing Antibody in Chimpanzees

During the first 17 months of the immunization trial, serum-neutralizing antibodies against HIVIIIB were always undetectable in syncytium inhibition assay and were  $\leq 1:45$  in reverse transcriptase inhibition assay. However, following immunization of animals 1045 and 1070 at month 17 with HIVMN Th-B peptide, anti-HIV neutralizing antibodies were seen in syncytium inhibition assay (Table 11).

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To determine why antibodies against HIVIIIIB Th-B peptides did not neutralize HIVIIIIB in vitro during the first 17 months of immunization, sera from the early peak anti-HIVIIIIB peptide antibody responses (month 6) were assayed for reactivity to the individual epitopes of the Th-B peptides. It was found that at the time of initial titers of anti-Th-B peptide responses, most of the antibody reactivity in sera from animals 884 and 1028 was indeed directed to the primary amino acid sequence of the neutralizing V3 loop region defined by the peptide (TRKSIRIQRGPR) (Table 12,). These data indicate that antibodies made by chimpanzee nos. 884 and 1028 at 7 months after immunization with the HIVIIIIB Th-B HIV env peptides did not recognize the appropriate secondary V3 loop structure(s) necessary for neutralizing HIVIIIIB, although the animals did make antibody responses to the correct primary amino acid sequences of the neutralizing V3 B cell determinant of HIVIIIIB gp120.

#### Example 9

Regarding the induction of tolerance, additional clinical syndromes that might be treated using Fusion domain or Fusion domain-like peptides synthesized N- or C-terminal to an otherwise immunogenic antigen is in hypersensitivity to bee or wasp venom antigens and hypersensitivity to plant or animal allergens. The nucleotide and amino acid sequences of a number of allergens have now been synthesized, and those regions of the allergen proteins that induce IgE antibodies or T

helper cell responses that help to induce IgE  
 responses are being mapped. Thus the primary  
 structure of grass pollen (Silvanovich et al J.  
 Biol. Chem. 266:1204-1220, 1991; Griffith et al  
 5 FEBS Letters 279:210-215, 1991; Perez et al J.  
 Biol. Chem. 265:16210-16215, 1990; Singh et al  
 Proc. Natl. Acad. Sci. USA 88:1384-1388, 1991),  
 mite allergens (Tovey et al J. Exp. Med. 170:1457-  
 1462, 1989; Yasel et al J. Immunol. 148:738-745  
 10 1992; Chua et al J. Exp. Med. 167:175-182, 1988;  
 Chua et al Int. Arch. Allergy Appl. Immunol.  
 91:118-123, 1990), hornet venom (Fang, et al Proc.  
 Natl Acad. Sci., USA 85:895-899, 1988), and tree  
 pollen (Ebner et al J. Immunology 150:1047-1054,  
 15 1993; Jarolim et al Int. Arch. Allergy Appl.  
 Immunol. 90:54-60, 1989; Valenta et al Science  
 253:557-560, 1991). For some of these allergen  
 proteins, T cell epitopes have been mapped (Ebner  
 et al J. Immunology 150:1047-1045, 1993) while for  
 20 others, likely T cell sites and hydrophilic B cell  
 determinants can be predicted using computer  
 algorithms (Kyte and Doolittle J. Mol. Biol.  
 157:105-132, 1982; Rothbard and Taylor EMBO J.  
 7:93, 1988; Margalit et al J. Immunol. 138:2213,  
 25 1987) and tested by synthesizing peptides and  
 injecting animals, or by reacting patient serum  
 antibodies or peripheral blood T cells with  
 synthesized peptide in in vivo assays. Once

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identified, T and B cell epitopes of bee, wasp or other allergens can be synthesized with a F domain or F-domain-like peptide N- or C-terminal to the allergenic T or B cell peptide, and the F-Allergen epitope hybrid peptide used to inject into patients that are sensitive to the allergen epitope. By this method, a patient can be made tolerant to the allergen epitope in the same manner as chimpanzees were made tolerant to T1-SP10 HIV env peptides by immunizing them with F-T1-SP10(A) peptide (Haynes et al J. Exp. Med. in press, 1993). Thus, in addition to treating autoimmune disease, F-derivatizing allergen T or B cell immunogenic peptides could product tolerogenic peptides for the treatment of allergic diseases.

A new technology has been developed whereby injection in vivo of cDNAs with a powerful promoter and encoding immunogenic peptides or proteins has been found to promote internalization and expression of cDNAs in host cells (Wolff et al Science 247:1465, 1990). Thus, the above strategy could be performed whereby cDNAs encoding F-derivatized peptides of autoantigens and/or allergens are injected instead of the peptides themselves, thus having the same effect as immunizing with peptides themselves. Moreover, the F-derivatized peptides and proteins could be

produced by recombinant DNA techniques instead of peptide synthesis of peptide synthesis and the same type of tolerizing immunogen obtained.

Another use of F domain- or F-like domain

5 derivatization of peptides and proteins is to confer upon the derivatized peptide or protein the ability to bind to the cell membrane and enter the cell. The fusion domain or a fusion-like domain could be conjugated to an RNA or DNA molecule as

10 well as a protein to promote entry into cells. The ability of a molecule to enter the cells is important for many molecules to act therapeutically, and can be overcome by addition of the F domain or an F-like domain to the

15 molecule that one wanted to get inside or cells. For example, a powerful inhibitor of cell activation would be a peptide, RNA or DNA species of molecule that competitively bound to an intracellular molecule necessary for cell

20 activation, but the peptide, RNA or DNA molecule itself did not activate or serve the normal function of the physiologic ligand that it was designed to mimic. Examples of peptide, RNA or DNA molecules that might inhibit cell activation

25 would be molecules that bound to intracellular tyrosine kinases, tyrosine phosphatases, protein Kinase C enzymes or G proteins, just to name a few examples. However, for peptide, RNA or DNA

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inhibitory ligands to function as cell regulatory  
agents when administered as therapeutic agents,  
they must readily bind the cell without killing  
the cell, and be able to enter the cell and  
5 function intracellularly. It has been shown that  
F-derivatization of the T1-SP10IIIB(A) peptide  
with the HIV gp41 F domain promotes enhanced  
binding (Table 13) and entry (Table 14) of the  
derivatized T12-SP10IIIB(A) peptide into human B  
10 cells. This ability to promote entry of  
derivatized molecules into the inside of cells  
represents a novel drug delivery system with  
potential uses for delivering virtually any type  
of molecule (RNA, DNA, protein) inside cells for  
15 the desired therapeutic effect. For example, F-  
derivatized proteins of HIV regulatory proteins  
that might bind to viral RNA but not promote  
transcription of RNA thus preventing normal  
binding of HIV transcription factors might be used  
20 to treat HIV infections in vivo.

\* \* \* \* \*

All publications mentioned hereinabove  
are hereby incorporated in their entirety by  
reference.

25 While the foregoing invention has been  
described in some detail for purposes of clarity

and understanding, it will be appreciated by one skilled in the art from a reading of this disclosure that various changes in form and detail can be made without departing from the true scope of the invention and appended claims.

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